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Wong, Swee Chong

2016

Wong , S C , Oksanen , A , Mattila , A L K , Lehtonen , R , Niitepöld , K & Hanski , I A 2016 ,
' Effects of ambient and preceding temperatures and metabolic genes on flight metabolism in
the Glanville fritillary butterfly ' , Journal of Insect Physiology , vol. 85 , pp. 23-31 . <https://doi.org/10.1016/j.jinsphys.2015.11.015>

<http://hdl.handle.net/10138/217888>

<https://doi.org/10.1016/j.jinsphys.2015.11.015>

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Effects of ambient and preceding temperatures and metabolic genes on flight metabolism in the Glanville fritillary butterfly



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ARTICLE INFO

Article history:

Received 14 September 2015

Received in revised form 26 November 2015

Accepted 30 November 2015

Available online 3 December 2015

Keywords:

Flight metabolic rate

Temperature treatment

Pgi

Tpi

Temperature–genotype interaction

Flight muscle maturation

ABSTRACT

Flight is essential for foraging, mate searching and dispersal in many insects, but flight metabolism in ectotherms is strongly constrained by temperature. Thermal conditions vary greatly in natural populations and may hence restrict fitness-related activities. Working on the Glanville fritillary butterfly (*Melitaea cinxia*), we studied the effects of temperature experienced during the first 2 days of adult life on flight metabolism, genetic associations between flight metabolic rate and variation in candidate metabolic genes, and genotype–temperature interactions. The maximal flight performance was reduced by 17% by 2 days of low ambient temperature (15 °C) prior to the flight trial, mimicking conditions that butterflies commonly encounter in nature. A SNP in *phosphoglucose isomerase* (*Pgi*) had a significant association on flight metabolic rate in males and a SNP in *triosephosphate isomerase* (*Tpi*) was significantly associated with flight metabolic rate in females. In the *Pgi* SNP, AC heterozygotes had higher flight metabolic rate than AA homozygotes following low preceding temperature, but the trend was reversed following high preceding temperature, consistent with previous results on genotype–temperature interaction for this SNP. We suggest that these results on 2-day old butterflies reflect thermal effect on the maturation of flight muscles. These results highlight the consequences of variation in thermal conditions on the time scale of days, and they contribute to a better understanding of the complex dynamics of flight metabolism and flight-related activities under conditions that are relevant for natural populations living under variable thermal conditions.

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1. Introduction

Temperature influences all biological processes. Ectothermic animals in particular are greatly affected by thermal conditions, as both high and low temperatures constrain activity and reduce fitness. At geographical scale, adaptation to the thermal environment is exemplified by *Drosophila melanogaster* in North America and Australia, where there are distinct temperature-related latitudinal clines in several life history and morphological traits as well as in molecular variation in central metabolic genes (Hoffmann and Weeks, 2007; Sezgin et al., 2004; Reinhardt et al., 2014; Cogni et al., 2015). At a smaller spatial scale, the willow beetle *Chrysomela aeneicollis* that inhabits the Sierra Nevada Mountains in California shows temperature-associated geographic variation in the glycolytic allozyme *phosphoglucose isomerase* (PGI) (Dahlhoff and

Rank, 2000). In this case, variation in allele frequencies reflects genotypic differences in thermal stability at the enzyme level and heat and cold tolerance at the individual level (Dahlhoff and Rank, 2007). Moreover, the thermally sensitive genotype shows higher heat shock protein *Hsp70* expression levels (Neargarder et al., 2003). In North American *Colias* butterflies, allozyme variation in PGI is associated with enzyme kinetics and thermal stability, which in turn has been shown to correlate with flight activity, mating success and reproductive performance (Watt, 1977; Watt et al., 2003). In butterflies in general, flight performance is critical for fitness in the wild, and flight activity is highly dependent on temperature.

The flight muscles of temperate butterflies operate best within a narrow range of temperatures, which is well above the typical ambient air temperatures (Kohane and Watt, 1999; Kingsolver, 1983). Work on the Eurasian Glanville fritillary butterfly (*Melitaea cinxia*) has shown that within the favorable range of temperatures, molecular variation in the *Pgi* locus is associated with variation in

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flight metabolic rate (FMR) and flight performance in the field (Niitepöld, 2010; Orsini et al., 2009). A single nucleotide polymorphism (SNP) in the *Pgi* locus interacts with ambient temperature in affecting both FMR and mobility: one genotype shows higher activity in lower temperatures, while the other one performs better in higher temperatures (Niitepöld et al., 2009).

Flight metabolic rate is a fundamentally important process for butterflies and many other insects, but it remains an understudied trait and we lack fundamental knowledge of its dynamics and the environmental conditions affecting it. For instance, though the effects of ambient temperature on butterfly activity and performance have been well documented, we know much less about the effects of thermal conditions experienced by individuals prior to particular flight activity. Temperature is never constant on the scale of days, and many insects are short-lived, which means that thermal conditions experienced during early life can potentially have a large effect on fitness. A previous study on the Glanville fritillary indicated that the isoform composition of *tropoin t*, an important muscle gene associated with performance, changes significantly in early life, presumably reflecting gradual maturation of flight muscles following adult eclosion (Marden et al., 2008). Thermal conditions may influence the rate of maturation, and so may genetic variation in metabolic genes, which suggests that flight metabolism of young butterflies may be sensitive to the thermal conditions they have experienced earlier in their life, with possible interactions with genetic variation.

In the present study, we investigated how the thermal conditions during a few days following adult emergence affect flight performance in the Glanville fritillary butterfly, a species that inhabits fragmented habitats and whose ecological dynamics are therefore highly dependent on flight performance (Hanski, 1999, 2011). We focus on three questions. First, we tested whether thermal conditions in the first two days after emergence, when the butterfly is fully formed but still undergoes physiological maturation, have an effect on flight metabolic rate (FMR), a quantitative measure of flight performance. Second, we conducted an association study on SNPs in candidate metabolic genes using mixed model regression. Third, we examined whether these candidate SNPs interact with previously experienced temperatures versus ambient (current) measurement temperature on affecting FMR. The results demonstrate significant effects of both current and past temperatures on FMR. Among the genetic associations, we found a significant association of the SNP in *Pgi* but also a novel association between FMR and a SNP in the gene *triosephosphate isomerase* (*Tpi*), which is located on the sex chromosome. To test the generality of this result, the SNP was genotyped in two additional data sets from previous experiments, and was found to have a significant result in one of them.

2. Materials and methods

2.1. Study materials

The primary dataset, hereafter referred to as DS1, consists of the offspring of butterflies studied in an outdoor population cage experiment at the Lammi biological station in Finland in 2008 (Klemme and Hanski, 2009). The parents of these offspring had been sampled as 5th instar larvae from the field, across the Åland Islands (50 by 70 km) in SW Finland, in the autumn of 2007. For the present experiment, we sampled the offspring from 15 mating. The offspring were reared to the 5th instar in the laboratory and maintained in diapause at 3 °C over winter. In spring 2009, post-diapause larvae were reared into adult butterflies in controlled conditions (12 h/12 h L/D, 28 °C/8 °C), fed with the leaves of the host plant *Plantago lanceolata*.

Two additional datasets, called DS2 and DS3, were used to validate the novel discovery from DS1, and these data were originated from the study of Kvist et al. (2015) and Mattila (2015). Both DS2 and DS3 consist of apparently unrelated individuals originating from 31 and 87 larval family groups (full sibs) sampled across the Åland Islands in 2011 and 2010, respectively. Flight metabolic rate was measured in the same way as in DS1 (below), but there was no temperature treatment. The samples and rearing conditions for DS1, DS2 and DS3 are described in [Supplementary Table S1 in the Supplementary Information](#).

2.2. Temperature treatments and the measurement of metabolic rate

Adult butterflies from DS1 were marked individually after eclosion and assigned into three thermal treatments: 15, 24 and 35 °C. The treatment temperatures were chosen based on the activity level of butterflies at different temperatures. At 15 °C, butterflies are inactive, they do not move nor feed and often rest with their wings closed. At 35 °C, butterflies are very active and constantly fly around in the cage (diameter 15 cm, height 30 cm) in the laboratory. The intermediate temperature, 24 °C, is favorable for maintaining butterflies in the laboratory. Each group was kept in their designated temperature for 12 h from 0800 until 2000 during two days, and they all spent the night (2000–0800) at 8 °C. Butterflies were provided with water in a sponge but no sugar solution.

The flight metabolic rate of 51 females and 89 males was measured as CO₂ emission rate during flight on the third day following eclosion (see Niitepöld et al. (2009) for details on the method). Half of the individuals were measured at 30 °C and the other half at 35 °C. These two temperatures were selected as we expected heterozygous *Pgi* AC individuals to show higher performance in lower temperatures and homozygous AA individuals to perform better in higher temperatures (Niitepöld et al., 2009). Butterflies were encouraged to fly for 10 min as continuously as possible by gently shaking and tapping the one-liter respirometry chamber whenever the individual landed. The chamber was kept under a light source emitting both visible and UV light. We extracted two variables from the data to describe metabolic performance during flight: the highest rate of CO₂ production (FMR_{peak}), which typically occurs within a few min from the beginning of the experiment, and the total (integrated) volume of CO₂ (FMR_{int}) produced during the 10-min experiment. FMR_{peak} reflects maximal metabolic performance whereas FMR_{int} reflects a combination of endurance and behavior. The experimental protocols for DS2 and DS3 were similar to that for DS1 except for the duration of the experiment and lack of the temperature treatment prior to the measurement of metabolic rate. In DS2, metabolic rate was measured for 16 females and 11 males for 15 min, while in DS3 it was measured for 36 females and 51 males for 7 min. Measurement temperature was 30 °C in both DS2 and DS3. Further details on the three datasets are given in the [Supplementary Table S1](#).

2.3. Selection of candidate genes and genotyping

In DS1, 18 SNPs from 12 Expressed Sequence Tag (EST) sequences and 3 SNPs from the *phosphoglucose isomerase* (*Pgi*) partial messenger RNA (mRNA) sequence were selected for genotyping ([Supplementary Table S2](#)). These sequences were extracted from transcriptome data (Ahola et al., 2015) where complementary DNA (cDNA) from individuals originating from the Åland Islands in Finland, China and France were sequenced with 454 pyrosequencing (Roche Diagnostics, US; GS FLX). The resulting sequences were annotated with PANNZER (v 1.0) annotation tool (Radivojac et al., 2013). Gene set enrichment analyses (GSEA) were performed for pairs of populations to explore the enrichment of gene ontologies (GO). Significant results from GSEA were used to compile the initial

set of genes. As our aim was to determine genetic associations with flight metabolic rate, we selected mostly genes of central metabolic pathways as candidate genes, such as genes from the glycolysis pathway (*phosphoglucose isomerase*, *triosephosphate isomerase*), the pentose phosphate pathway (*glucose-6-phosphate dehydrogenase*), and the oxygen binding pathway (*cytochrome P450 337*). Stress response genes (*Heatshock protein 70*, *c-Jun N-terminal protein kinase 1*) were also included as candidate genes based on previous studies (Czaja, 2010; Karl et al., 2008). Finally, several other genes were selected based on significant results in previous gene expression studies on life-history traits in the Glanville fritillary (*succinate dehydrogenase complex subunit D*, *flightin*, *peripheral-type benzodiazepine receptor*, *troponin t*) (Klepsatel and Flatt, 2011; Marden et al., 2008; Wheat et al., 2011) (Supplementary Table S2). Candidate SNPs in these genes with their flanking sequences were extracted from the corresponding EST for genotyping, which was performed with Sequenom iPLEX Gold (Sequenom Inc. CA, USA) chemistry in a single multiplex well at the Institute for Molecular Medicine Finland (FIMM, Helsinki). Primers and probes for genotyping were designed using the MassARRAY Assay Designer Program (2006).

A SNP in the gene *triosephosphate isomerase* (*Tpi*) was significantly associated with female FMR in DS1. As this association has not been reported previously, we genotyped females from two other experiments on FMR to validate this finding. In DS2, the SNP was identified from RNA-seq data in Kvist et al. (2015). RNA was extracted from thorax and sequenced using next generation RNA sequencing technology implemented in Illumina HiSeq2000 and HiScanSQ (Illumina Inc., San Diego, CA, USA). SNP and genotype calling were performed using SAMtools ver 0.1.18 (Li et al., 2009) in two steps. The first step involved piling up the relevant input file in the .bam format in relation to the reference genome (Ahola et al., 2014). The output of this “pileup” step was piped to bcftools to create a single bcf file. In the second step, variant calling was performed by first viewing the bcf file via bcftools, and the output was piped to vcftools for variant calling (Full command in the Supplementary material). In DS3, we used Sanger sequencing to determine the genotypes for the SNP, which is located in the 3' UTR region of *Tpi*.

2.4. Statistical analyses

Mixed model multiple regression was used to test for the effects of body mass (pupal weight), preceding temperature and measurement temperature on flight metabolic rates (FMR_{int} and FMR_{peak}), with brood as a random factor. As sex-dependent genetic effects occur in insects (Le Goff et al., 2006), the data were analyzed separately for males and females to facilitate the interpretation of the results. The statistical analyses were performed using R (x64 ver 3.1.0) (R Core Team, 2014).

For dataset DS1, association studies were implemented with mixed model regression using the “nlme” package in R (Pinheiro et al., 2014). To control for the relatedness of individuals, brood was used as a random variable in the model. Body mass was treated as a covariate, while preceding and measurement temperatures were treated as factors. To determine the significance of each explanatory variable in models for FMR, Anova function in the R package “car” (Fox and Weisberg, 2011) was used to obtain *P* values from the Wald Chi-square test. The *P* values were adjusted for multiple testing via false discovery rate (FDR) (Benjamini and Hochberg, 1995). We used the Lund test (Lund, 1975) for outlier detection to exclude significant outliers before the regression model.

In the validation datasets (DS2 and DS3), apparently unrelated samples were collected from the wild, but additional measures were taken to account for possible population stratification. In

DS2, genome-wide autosomal SNPs were extracted from RNA-seq data to construct a matrix of relationships of the individuals. SNPs were filtered by minor allele frequency (MAF > 0.2), call rate (CR > 0.9) and linkage disequilibrium (LD) scores ($R^2 < 0.5$, higher R^2 indicates that SNPs are in LD), yielding 105 SNPs for the estimation of the relationship matrix to control for population stratification. Dataset DS2 was analyzed using mixed model regression implemented in Tassel software package ver 5.1.0 (Bradbury et al., 2007; Zhang et al., 2010). In DS3, individuals were sampled from different habitat patches across the large 4000 patch network in the Åland Islands (Hanski, 2011). As there was no genome-wide SNP data available in DS3 to construct the relationship matrix, we used sub-networks into which the 4000 patch network has been subdivided as a parameter to control population stratification in a linear model implemented in PLINK (Purcell et al., 2007). The properties of each dataset and the methods of their analysis are summarized in Supplementary Table S1. We report in the Supplementary material power analyses for the statistical tests.

3. Results

We excluded 3 of the 21 SNPs as they showed no variation (Supplementary Table S2). Two pairs of SNPs were completely linked and hence only one SNP from each pair was retained, resulting in 16 SNPs from 11 genes (detailed characterization of SNPs are depicted in Supplementary Tables S2 and S4). Below, we present the results for a series of increasingly complex models. We first analyze the effects of preceding temperature treatments and measurement temperatures on flight metabolic rates (FMR_{int} and FMR_{peak}), then test for the associations of the SNPs, and finally analyze possible interactions between the SNPs and the temperature before and during the measurement of FMR.

3.1. Effects of preceding temperature and measurement temperature on FMR

In males, body mass, the 2-day temperature treatment and the measurement temperature all had significant effects on FMR_{peak} (Table 1). In the case of FMR_{int} , the result was qualitatively the same, but the effect of body mass was not quite significant. The effect size of the preceding temperature treatment was large, with 17% greater values of FMR_{peak} at 35 °C than at 15 °C. In females, body mass and preceding temperature treatment had significant effects on FMR_{int} , while the effect of preceding temperature was nearly significant on FMR_{peak} (Table 1). According to the model, changing the preceding temperature from 15 to 35 °C increased FMR_{peak} by 16% in females. The effects of the explanatory variables on FMR_{peak} are illustrated in Fig. 1.

Table 1

The effects of body mass (pupal weight), 2-day temperature treatment preceding the measurement, and the measurement temperature on the integrated and peak flight metabolic rates in males and females. Significant results are highlighted in bold.

Sex	Trait	Explanatory variable	χ^2	df	$Pr(>\chi^2)$
Male	FMR_{int}	Pupal weight	3.273	1	0.070
		Preceding temp	5.654	1	0.017
		Measurement temp	7.606	1	0.006
	FMR_{peak}	Pupal weight	38.670	1	5.02E–10
		Preceding temp	6.161	1	0.013
		Measurement temp	19.313	1	1.11E–05
Female	FMR_{int}	Pupal weight	6.125	1	0.013
		Preceding temp	8.670	1	0.003
		Measurement temp	0.014	1	0.906
	FMR_{peak}	Pupal weight	21.247	1	4.04E–06
		Preceding temp	2.784	1	0.095
		Measurement temp	2.990	1	0.084

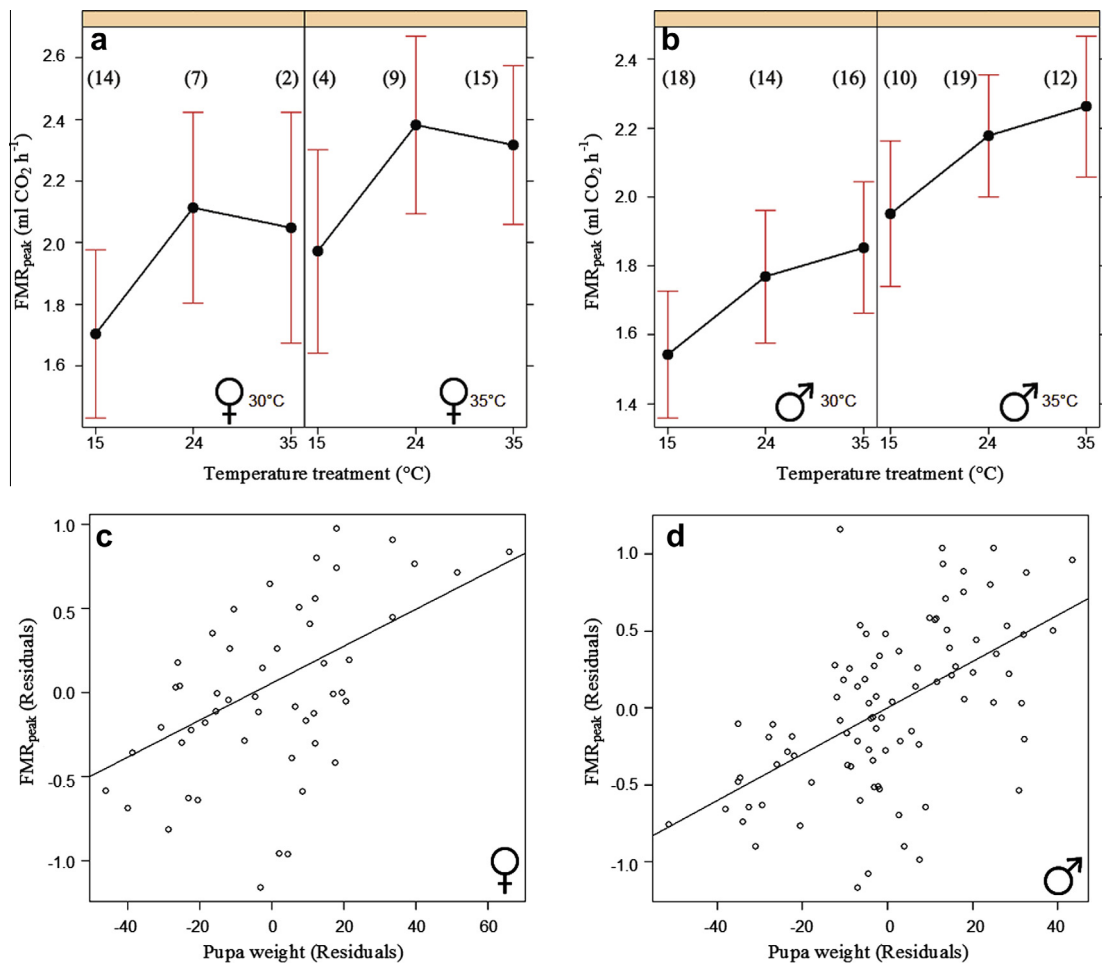


Fig. 1. The effect plots with 95% confidence intervals for three 2-day temperature treatments prior to the measurement and two measurement temperatures on FMR_{peak} in (a) females and (b) males. Partial regression plots of FMR_{peak} against body mass are shown for (c) females and (d) males, controlling for the preceding temperature treatment and measurement temperature. Sample size is shown in parentheses close to the data point.

3.2. Association between SNPs and FMR

Table 2 gives the results for the SNPs with significant association with metabolic rates (false discovery rate < 0.05). These associations were detected in SNPs from two metabolic genes, *phosphoglucose isomerase* (*Pgi*) and *triosephosphate isomerase* (*Tpi*). In the case of the *Pgi* SNP, *pgi:c.331A>C*, the association with integrated flight metabolic rate (FMR_{int}) was significant in males, while the SNP from *Tpi*, *tpi:1250A>G*, was significantly associated with both FMR_{peak} and FMR_{int} in females. Fig. 2 shows the results for FMR_{int} and *pgi:c.331A>C* in males and for *tpi:1250A>G* in females. The full association results for all the SNPs are presented in Supplementary Table S3.

3.3. Interaction between SNPs and temperature on FMR

We next analyzed possible interactions between SNPs and the preceding temperature treatment and the measurement temperature. For the measurement temperature, none of the interactions were significant, but there was one significant interaction between preceding temperature and a SNP from *Heatshock protein 70*, *hsp_4:106A>G*, in females (FMR_{int} $P = 0.0026$, FDR = 0.042; FMR_{peak} $P = 0.0009$, FDR = 0.014). One additional copy of the A-allele increased FMR_{peak} at the highest temperature treatment, in clear contrast to the trend in the lower temperature treatments (Fig. 3a).

Table 3 shows trait value differences of *pgi:c.331A>C* genotypes on FMR in the three preceding temperature treatments. The same

Table 2
Associations of SNPs in two metabolic genes with integrated and peak flight metabolic rates. Significant associations are highlighted in bold.

Sex	Trait	SNP	Accession	Allele	Effect	P	FDR
Male	FMR_{int}	<i>pgi:c.331A>C</i>	EU888473.1	C	0.028	0.0017	0.0265
		<i>tpi:1250A>G</i>	KJ803028	G	0.004	0.7257	0.9213
	FMR_{peak}	<i>pgi:c.331A>C</i>	EU888473.1	C	0.179	0.0559	0.1774
		<i>tpi:1250A>G</i>	KJ803028	G	−0.032	0.6783	0.6783
Female	FMR_{int}	<i>pgi:c.331A>C</i>	EU888473.1	C	−0.006	0.6236	0.9070
		<i>tpi:1250A>G</i>	KJ803028	G	0.070	0.0008	0.0120
	FMR_{peak}	<i>pgi:c.331A>C</i>	EU888473.1	C	−0.137	0.2259	0.6824
		<i>tpi:1250A>G</i>	KJ803028	G	0.490	0.0024	0.0381

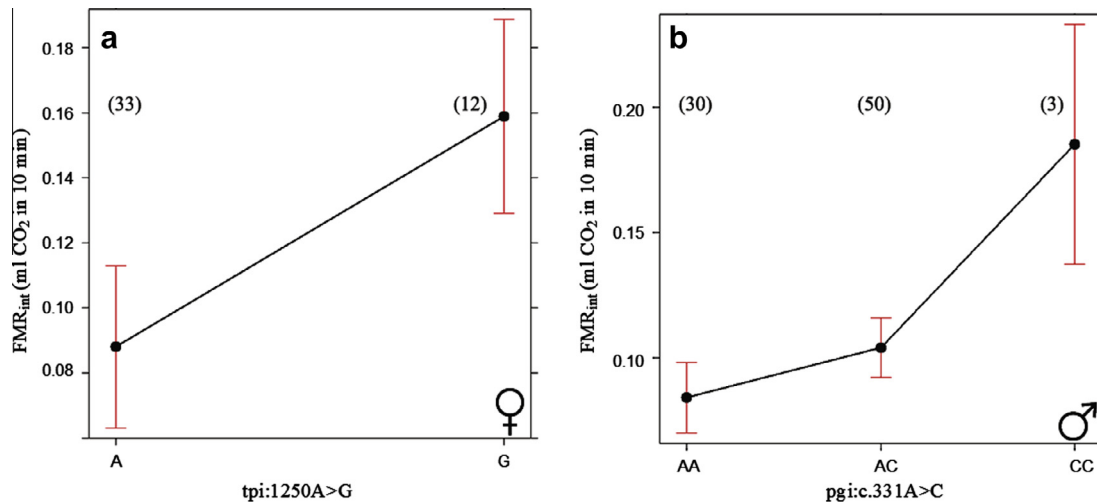


Fig. 2. The genotypic plot with 95% confidence intervals of SNP (a) tpi:1250A>G in females and (b) pgi:c:331A>C in males on FMR_{int}. SNP tpi:1250A>G is Z chromosome-linked, hence there is only one copy of the allele in females. Sample size is shown in parentheses close to the data point.

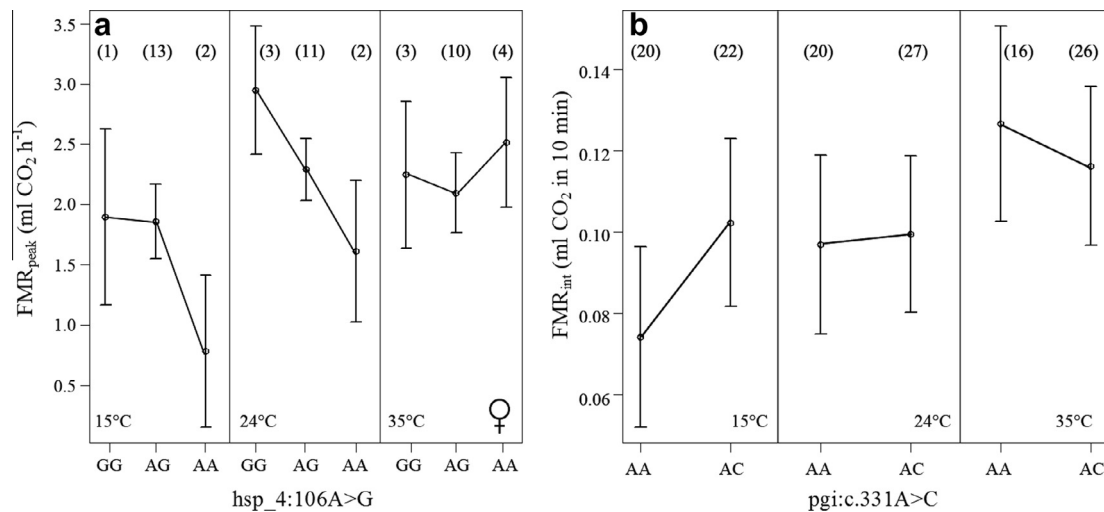


Fig. 3. (a) The genotypic plot with 95% confidence intervals at three different preceding temperature treatments for the SNP hsp_4:106A>G on FMR_{peak} in females, and (b) for SNP pgi:c:331A>C in pooled data for males and females. Sample size is shown in parentheses close to the data point.

Table 3

Genotypic differences of pgi:c:331A>C between the AC and AA individuals in the different preceding temperature treatments. Positive values indicate higher values of AC than AA individuals. The differences are significant for FMR_{int} ($P=0.017$) and FMR_{peak} ($P=0.016$) in males at 15 °C, which are highlighted in bold. For further analysis see the text and Table 4.

	Males Effect (%)			Females Effect (%)		
Preceding temp (°C)	15	24	35	15	24	35
N (AC, AA)	14, 13	19, 12	17, 10	8, 7	8, 8	9, 6
FMR _{int}	56.3	19.2	14.8	7.5	2.7	-7.1
FMR _{peak}	22.5	16.6	-4.7	13.3	-17.1	-9.2

pattern is seen in both sexes and in both FMR_{peak} and FMR_{int}: the AC heterozygotes had higher FMR than the AA homozygotes in the lowest preceding temperature treatment, but vice versa at higher preceding temperatures (Table 3). As the pattern was the same in the two sexes, we analyzed the pooled data for FMR_{peak}, including temperature treatment prior to flight, measurement temperature, sex and the SNP pgi:c:331A>C as well as their interactions as explanatory factors. The best model based on Akaike Information Criterion (AIC) included all the main factors as well as interactions between the SNP and sex and the SNP and

the preceding temperature (Table 4). The latter interaction gives statistical support to the pattern in Table 3.

3.4. Validation of the association between tpi:1250A>G and FMR

The association between the SNP tpi:1250A>G and FMR in females has not been reported previously in the literature, and

Table 4

Models for the peak and integrated flight metabolic rate involving the SNP pgi:c:331A>C using pooled data for males and females. Significant results are highlighted in bold.

Explanatory variable	FMR _{int}			FMR _{peak}		
	χ^2	df	Pr(> χ^2)	χ^2	df	Pr(> χ^2)
Pupal weight	8.1018	1	0.00442	55.2452	1	1.06E-13
Preceding temp	16.1818	1	5.75E-05	12.115	1	0.0005
Measurement temp	8.1117	1	0.0044	25.6542	1	4.08E-07
Sex	0.596	1	0.44012	0.3529	1	0.5525
pgi:c:331A>C	11.9132	1	0.00056	9.9502	1	0.00161
Sex: pgi:c:331A>C	5.2068	1	0.0225	4.3131	1	0.03782
Preceding temp: pgi:c:331A>C	5.4732	1	0.01931	5.315	1	0.02114

Table 5

Association of the SNP tpi:1250A>G with integrated and peak flight metabolic rates in females in datasets DS2 and DS3. Significant result is highlighted in bold.

Trait	Dataset DS2		Dataset DS3	
	F	P	T	P
FMR _{int}	6.258	0.028	−1.378	0.179
FMR _{peak}	1.641	0.224	−1.958	0.060

hence we validated this result with material from two additional datasets, DS2 and DS3. In DS2, genotypes were available for 16 females. One significant outlier was removed from the dataset before the analysis. The SNP tpi:1250A>G was significantly associated with FMR_{int} ($P = 0.028$) but not with FMR_{peak} (Table 5). In dataset DS3, there was no significant association with either measure of flight metabolic rate (Table 5).

4. Discussion

4.1. Effects of body mass and preceding temperature on FMR

Previous studies on the Glanville fritillary (Niitepöld et al., 2009) and other insects (Harrison and Fewell, 2002; Harrison et al., 1996) have demonstrated an effect of ambient temperature on flight metabolic rate when a wide range of temperatures has been studied. We measured flight metabolic rate at two temperatures, 30 and 35 °C, which are both favorable for flight, as butterfly flight muscles perform best when their temperature is between 30 and 40 °C (Kohane and Watt, 1999; Watt, 1968). For these two temperatures, we found a significant effect of measurement temperature on flight metabolism in males, especially on FMR_{peak}, but no significant effect in females. In the experiment, butterflies were moved into the respirometry chamber from the temperature of 22 °C ca. 25 min before the measurement. As female butterflies have larger body mass than males (19% difference in pupal weight in the present material), males can be expected to be more sensitive to temperature changes (Gilchrist, 1990). In the field, male butterflies spend shorter periods of time basking before flight (Gilchrist, 1990), as they can increase body temperature faster than females. In an experiment on the Glanville fritillary, thoracic temperature during take-off was positively correlated with ambient temperature in males but not in females (Mattila, 2015). These results suggest that body temperature responds more slowly to short-term changes in ambient temperature in females than in males, which may explain the sex difference in the effect of measurement temperature in our results.

Thermal conditions experienced during the first two days following adult eclosion had a significant effect on flight metabolic rate. In females, FMR_{int} was 53% greater in individuals that had spent two days at 35 °C compared to 15 °C, and in males the difference was 28%. The corresponding figures for FMR_{peak} were 16% in females and 17% in males. The temperatures used in our treatments correspond to realistic thermal conditions experienced by butterflies in temperate climates. Butterflies can elevate their body temperature well above the ambient air temperature by basking in the sun, but under overcast conditions the body temperatures follow closely the ambient air temperature (Kingsolver, 1983).

We suggest that the mechanism explaining our results could be related to slower flight muscle maturation in lower temperatures. The maturation process of insect flight muscles involves changes in the abundance of alternatively spliced isoforms of the regulatory protein *troponin t*, as demonstrated for dragonflies (Fitzhugh and Marden, 1997; Marden et al., 1999), honeybees (Schippers et al., 2006), and Lepidoptera (Marden et al., 2008). The activity of several metabolic enzymes increases during the first days of life in

honeybees, and the flight metabolic rate of honeybees increases in early life, and also later in life, when workers undergo a behavioral change from staying in the hive to foraging in the field (Schippers et al., 2010). In butterflies, flight metabolic rate has been shown to be low on the first day after emergence, after which it increases. Mated females of *Speyeria mormonia* reached their highest flight metabolic rate four days after emergence (Niitepöld et al., 2014). In unmated male Glanville fritillaries, flight metabolic rate increased from the age of one day to the age of three days, after which it stayed relatively stable (Niitepöld and Hanski, 2013). Physical activity may affect muscle maturation, as artificially increased wing loading and therefore increased need for muscle output changed the relative abundances of different *troponin t* isoforms in the flight muscles of the fall armyworm moth (*Spodoptera frugiperda*), and in the Glanville fritillary there was a correlation between voluntary activity and the *troponin t* isoform composition (Marden et al., 2008). Similarly, bumblebees initiate foraging flights while they still experience maturation at the enzyme activity level (Skandalis et al., 2011), supporting the notion that activity is an important driving force in the maturation process. In summary, these results suggest that conditions experienced during the first days after adult emergence may interact with the process of maturation. To test this hypothesis, further experiments on *troponin t* isoform relative abundances and gene expression under different thermal conditions and levels of physical activity are needed.

Long-term thermal conditions during the larval stage may have different consequences than thermal conditions experienced during adult maturation. Mattila and Hanski (2014) found that larval development in high temperature (daytime temperature 35 °C) resulted in significantly reduced flight metabolic rate in adults in comparison with larval development in moderate temperature (daytime temperature 28 °C). High temperature during larval development may lead to a stress response with consequences for adult flight performance. Moreover, Mattila and Hanski (2014) found that FMR_{int} was significantly heritable in moderate temperature conditions (28 °C) but not under thermally stressful conditions (35 °C), highlighting the role of thermal plasticity in FMR. In the present study, variation among individuals in FMR was greater in females than in males (based on the variance of the random variable in the analysis). Heritability estimates for DS1 based on variance component linkage analysis (Amos, 1994; Sham et al., 2002) indicated high heritability of integrated flight metabolic rate (male $h^2 = 0.44$, female $h^2 = 0.65$), similar to the values reported by Mattila and Hanski (2014), but not of peak flight metabolic rate (male $h^2 = 0$, female $h^2 = 0.15$). Thus the more plastic of the two traits, FMR_{peak}, with a significant negative effect of preceding temperature, showed no significant heritability.

4.2. Association of metabolic SNPs

Previous studies on the Glanville fritillary have reported significant associations between the SNP pgi:c.331A>C in the glycolytic gene *phosphoglucose isomerase* and several life history traits, such as life span (Klemme and Hanski, 2009; Saastamoinen et al., 2009; Niitepöld and Hanski, 2013), clutch size (Saastamoinen, 2007; Saastamoinen and Hanski, 2008) and flight metabolic rate (FMR_{peak}) (Niitepöld and Hanski, 2013; Niitepöld et al., 2009; Niitepöld, 2010). Here, we found a significant association between pgi:c.331A>C and FMR_{int} in males but not in females, in agreement with the result of Mattila (2015). Pooled results for both sexes showed a significant interaction between the SNP and the preceding temperature treatment: AC heterozygotes had higher flight metabolic rate than AA homozygotes following two days at low temperature, but vice versa in the high preceding temperature treatment. This interaction is qualitatively similar to the one

between the *Pgi* genotype and measurement temperature when a wider range of measurement temperatures were used (Niitepõld et al., 2009). Moreover, the same interaction between *pgi*: c.331A>C and ambient temperature has been observed in the distance moved in a given time by free-flying female butterflies, tracked by harmonic radar: AC females flew longer distances than AA females in low to moderate ambient temperatures but not in high temperatures (Niitepõld et al., 2009). The present results suggest that the association is present also in post-emergence maturation, which has significant consequences for the performance and fitness of butterflies in the field. As discussed above, thermal conditions following adult emergence may influence flight muscle maturation, and as AC heterozygotes function better in cool conditions, they may mature at a faster rate than AA homozygotes in low temperatures. In this scenario, cool periods can affect butterfly mobility in the field through a direct effect on body temperature and through an indirect effect on maturation, affecting flight capacity later in life.

Triosephosphate isomerase (Tpi) is located in the same metabolic pathway as *Pgi* (glycolysis). It encodes the enzyme catalyzing the reversible conversion between *dihydroxyacetone phosphate* and *D-glyceraldehyde 3-phosphate*. Several insect studies have shown significant associations of genetic variation in *Tpi* on individual performance. For example, in a study on hypoxia tolerance in *D. melanogaster*, Zhou et al. (2008) discovered seven polymorphic sites in *Tpi* with distinct allele frequency differences between control and hypoxia tolerant individuals, including one SNP at the 3'UTR region. Another study on *D. melanogaster* showed a latitudinal cline in allele frequency in *Tpi* (Oakeshott et al., 1984).

Here, we detected a novel association between *tpi*:1250A>G and FMR in females. As *Tpi* is located in the sex chromosome, there is only one copy in females, while two copies in males, and hence there are no heterozygotes in females. In datasets DS1 and DS2, female butterflies with the G-allele exhibited higher flight metabolic rate than females with the A-allele. There was no significant association in DS3, possibly due to differences in the butterfly material and the experimental conditions (Supplementary Table S1), or due to insufficient power to detect true effects (sample size = 35). In any case, it is encouraging that two of the three independent datasets showed the same significant association between *FMR_{int}* and *tpi*:1250A>G. *Tpi* is thus another polymorphic metabolic gene that deserves attention in studies of metabolic rate and e.g. dispersal rate in insects. The dissimilar associations in the two sexes with the two metabolic genes, *Pgi* and *Tpi*, in the same pathway, deserve further attention. The gender difference may be related to the fact that *Tpi* is located in the sex chromosome, but it may also be affected by differences in flight behavior and hence differences in metabolic demands in the two sexes (Niitepõld et al., 2011).

Finally, SNP *hsp_4*:106A>G from one of the *Heat shock protein 70 (Hsp70)* genes interacted significantly with the preceding temperature treatment in influencing *FMR_{int}* and *FMR_{peak}* in females. An additional copy of the A-allele increased *FMR_{peak}* when the butterfly had spent 2 days at the highest temperature of 35 °C, while it had an opposite effect when the butterfly had experienced the temperatures of 15 and 24 °C (Fig. 3a). Heat shock proteins act as molecular chaperones protecting against cellular damage, and their expression is induced by heat and other stressors (Sørensen et al., 2003). The significant role of *Hsp* expression in thermal stress resistance is well established (Feder and Hoffmann, 1999), and it has been studied in many insects. For instance, in the willow beetle *C. aeneicollis*, thermally less tolerant *Pgi* genotypes up-regulate *Hsp70* to a greater extent than the other genotypes (Dahlhoff and Rank, 2000; Rank et al., 2007). Such buffering of individual differences in thermal tolerance by *Hsp* expression has been suggested also by other studies (Rutherford, 2003). Another study on insects shows that *Hsp* expression may be involved in the regulation of

thermal plasticity (Hu et al., 2014). On the other hand, over-expression of *Hsp* is expected to have deleterious fitness consequences (Krebs and Feder, 1997; Sørensen et al., 2003). In a previous study on the Glanville fritillary, Luo et al. (2014) found significant differences in *Hsp70* expression between populations originating from dissimilar thermal environments. Our results suggest that different *Hsp70* genotypes perform dissimilarly in relation to the thermal conditions experienced in early life. The differences may be due to e.g. differential expression of the *Hsp70* gene or its isoforms in different genotypes, which could affect stress response and regulation of flight metabolism. The genotypic differences may be reversed or only become visible during thermal stress when the expression of *Hsp70* is induced. However, due to small sample size in each interaction category in the present study (Fig. 3a), we consider that these results are not conclusive, in spite of statistical significance. Nonetheless, the present results suggest an interesting mechanism involving *Hsp70* that has not been reported before.

5. Conclusion

The present results demonstrate that thermal conditions experienced during 2 days following adult emergence have significant effects on subsequent flight metabolic rate in the Glanville fritillary butterfly. We found that the maximal flight performance was strongly reduced, by 16% to 17%, by 2 days of low temperature (15 °C) prior to the measurement of the flight metabolic rate. This response may represent slow flight muscle maturation in low temperatures where activity is limited. Such effects of post-emergence thermal conditions highlight the complex dynamics of flight metabolism, dispersal and insect behavior under conditions that are relevant for natural populations, and they introduce an additional source of variation in flight metabolism that is easily missed in typical laboratory experiments. The SNP in *phosphoglucose isomerase* interacted significantly with previously experienced thermal conditions in affecting flight metabolism, which may contribute to the maintenance of polymorphism in these loci in natural populations. These results highlight significant consequences of variation in thermal conditions on the time scale of days on butterfly flight metabolism and dispersal. The present results also point to new questions concerning plasticity of insect flight muscle maturation and the consequences for performance and fitness in natural populations.

Conflict of interest

The authors declare that they have no conflict of interest.

Acknowledgements

We thank Annukka Ruokolainen for DNA extraction and genotyping assay preparation; Institute for Molecular Medicine Finland (FIMM) for Sequenom genotyping; Minna Taipale, Karolinska Institutet and FIMM for RNA-seq sequencing; Panu Somervuo for suggestions and help with RNA-seq processing pipelines; and Virpi Ahola for EST data processing. We thank two anonymous referees for helpful comments. This study was supported by a grant from the Integrative Life Science (ILS) doctoral program to WSC, and grants from the European Research Council (AdG Grant Number 232826) and the Academy of Finland (Finnish CoE Programme, Grant Numbers 256453 and 250444) to IH.

A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at <http://dx.doi.org/10.1016/j.jinsphys.2015.11.015>.

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